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 EP-A- 0 346 981
 US-A- 3 326 723
 CHEMICAL ABSTRACTS, vol. 112, no. 6, 5th
 February 1990, page 577, abstract no.44303m,
 Cohambus, Ohio, US; N. MINEO et al.:
 'Zinc-iron alloy plating process by using hydrogen gas diffusion electrode as anode'
- (8) References cited:
 CHEMICAL ABSTRACTS, vol. 104, no. 4, 27th
 January 1996, page 172, abstract no. 22301s.
 Columbus, Dhio, US; G.M. CDOK et el.: "Using
 fuel cells or anodedepolarization to reduce
 electrowinning energy consumption:
 DERWENTAMPIL, accession no. 87-296109,
 Derwent Publications Ltd, London, GB; &
 JP-A-62 208 SS4 (TANAKA KIKINZOKU
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Description

Background of the Invention

The present invention relates to a process for electro placing a zinc alloy.

In these days, as a surface treated steel having a soft plating surface, stretchability, good affinity for a paint coating and excellent rust prevention effect which require no heal treatment in a field of automobiles, construction materials, appliances and the like, steels which are electroplated with such an alloy as zinc-iron, zino-cobalt, zino-nickel and zino-manganese have attracted public attention. Zino-iron plating will be described.

There are two processes for electroplating of a zinc-iron alloy, one of which employs a plating both containing sulfates of zinc and iron while the other employs a plating bath containing chlorides of zinc and iron. In case of the sulfate bath, such problems exist that liquid resistance is high and the deposition efficiency on a cathode is low because the pH is as low as 1.0. On the other hand, in case of the chloride bath, such advantages exist that liquid resistance is lower than that of the sulfate bath, the deposition efficiency on a cathode is good and the appearance is smooth.

However, in case of the chloride bath, a chlorine gas which is harmful to an anode evolves.

in case of the process for electropiating the zinciron alloy in which a trianium electrode plated with iridium or platinum is employed as an anode, the life of the electrode is as short as several thousand hours because the oxygen evolution occurs on the iridium or pial inum-pialed arlode. The valency of the iron lon in the plating bath increases in spite of the supplement of the pletting solution because the fron ion is oxidized on the anode. For this reason, the plating solution in the plating bath is circulated to an electrolytic reduction apparatus equipped with an ion exchange membrane to reduce Fe 3º to Fe2. Even by this procedure, the zinc, the iron ion and the pH of the platting bath become unbalanced so that, because the plating thickness may change and part of the surface may become rough, large steel companies discharge about two ton/day of the plating solution that causes serious loss

in Chem. Abstr., vol 112, no. 6, 05.02.90, page 577, abstr. no. 44903m, N. Mineo and N. Furuya (an inventor of the present application) disclose the plating of a zinc-tron alloy on an Iron plate (cathode) by using a hydrogen depolarized gas diffusion anode, in a chloride zinc plating bath. Reportedly, chlorine was not generated and the surface appearance of the plated oathode was excellent. In the document DER-WENT/WPIL, accession number 87/296109 corresponding to Japanese application JP-A-62208554, a gas diffusion electrode composed of a laminated film

having a reaction layer and a gas diffusion layer bounded together for use in fuel cells, electrochemical reactions and as anodes for electroplating, is disclosed.

Since the steel electroplated with the zinc-iron elloy moves at high speed of 200 m/min, in the small plating bath, the anode cannot be used if the mechanical strength is insufficient. The anode may be scratched and sparks when the anode contacts the steel.

Summary of the Invention

An object of the present invention is to provide a process for electroplating a steel with a zinc alloy using a gas diffusion electrode apparatus which does not require a frequent discharge of the plating solution.

A further object of the invention is to provide a process wherein scratches and sparks are prevented.

The process for electroplating a zino-iron alloy according to the present invention for overcoming the above problems comprises disposing in an electrolytic cell a gas diffusion electrode apparatus comprising a plurality of gas diffusion electrodes which function as an anode and are formed by a gas diffusion layer and a reaction layer being attached together, and an electroconductive reinforcing frame to which the gas diffusion electrodes are supported; forming a zinc alloy plating bath on the reaction layer side of the electrolytic cell which functions as anode chamber; supplying electricity to the gas diffusion electrodes through the reinforcing frame while supplying a hydrogen gas to the electrolytic cell of the gas diffusion layer side; and electroplating a steel with the zinc alloy by passing the steel as a cathode material through the zinc alloy bath.

Since in the process for electroplating the zinc allay of this invention the gas diffusion electrode is emplayed as an anode and the hydrogen gas is supplied to the gas diffusion layer side of the gas diffusion electrode to make an anode reaction of the platting bath to a hydrogen oxidation reaction, no oxygen evolves on the anode in case of the sulfate bath and no chlorine gas evolves on the anode so that a smooth and gorgeous zinc alloy plating layer without burns can be formed on the steel passing through the plating bath. Further, ions in the plating bath such as iron, cobalt, manganese and the like are not oxidized on the anode, and even if they are axidized on the anode to be contaminated in the plating bath in the form of their ions, these ions are reduced by the gas diffusion electrode to lower their ionic valency so that the zinc ion and the pH of the plating bath are never unbalanced. Accordingly, the plating bath can be effectively employed without the discharge of the plating solution, and the plating thickness of the zino-iron alloy applied to the steel does not change and the surface does not partially become rough.

Since a plurality of the gas diffusion electrodes are supported in the electroconductive reinforcing frame employed in the process for electroplating such an alloy-as zinc-iron, zinc-cobalt, zinc-manganese and the like as mentioned before, the mechanical strength of the gas diffusion electrode functioning as an anode is elevated. Therefore, if the steel moves at high speed in the small plating bath, the gas diffusion electrode is never flapped nor scratched and does not generate a spark.

Brief Description of the Drawings

Figs. 1 a and b show a gas diffusion electrode employed in the process of the present invention, Fig. 1 a being a plan view and Fig. 1 b being an A-A sectional view of Fig. 1 s; and

Fig. 2 shows a zinc-atioy electroplating bath employing the gas diffusion electrode apparatus of Fig. 1.

Detailed Description of the Invention

In the gas diffusion electrode apparatus shown in Figs. 1a and 1b, 1 designates a rectangular gas diffusion electrode. The gas diffusion electrode 1 is formed by a gas diffusion layer 2 and a reaction layer 3 attached together, the gas diffusion layer 2 consisting of hydrophobic carbon black and polytetrafluoroethylens, and the reaction layer 3 consisting of hydrophilic carbon black carrying a platinum catalyst, hydrophobic carbon black and polytetraftuoroethylene. A plurality of the gas diffusion electrodes 1, 10 electrodes in this embodiment; are instailed and supported in an electroconductive reinforcing frame made of titanium, copper, sixiniess and the like, in a lattice reinforcing frame 4 made of titanium in this embodimeni. A profection member 5 is formed on the front surface of the reinforcing frame 4 at the reaction layer 3 side of the gas diffusion electrode 1.

Agas diffusion electrode apparatus 9 employing the thus constructed gas diffusion electrode was disposed as an anode in an electrolytic cell 10, and a chloride bath 12 was filled in the cell chamber 11 formed on the reaction layer 3 side of the gas diffusion electrode 1, which defined an anode chamber 13. A hydrogen gas was successively introduced into the cell chamber 13 on the gas diffusion layer 2 side of the gas diffusion electrode 1, from an inlet port 14 so as to successively supply the hydrogen gas to the gas diffusion layer 2 side of the gas diffusion electrode apparatus 9 and to simultaneously discharge the hydrogen gas from the gas diffusion electrode 1 through a discharge port 15 of the cell chamber 13. At the same time, the current was passed to the respective gas diffusion electrodes 1 through the reinforcing frame 4. Under these circumstances, a steel

foli 16, as a cathode, was passed through the plating bath 12 of the cell cliember 11 at a relatively high speed.

By using the gas diffusion electrode 1 as an anode while supplying the hydrogen gas to the gas diffusion layer 2 side of the gas diffusion electrode 1 so as to make the anode reaction of the chloride plating bath 12 to be a hydrogen exidation reaction, no chlorine gas was evolved on the anode. The zinciron alloy layer plated on the steel 16 was examined to have a smooth and gorgeous surface, without burns. When the steel was successively passed through the cell to effect the electroplating of the zinoiron alloy, no iron lon was oxidized on the anode from Fe 2* to Fe 2*. Even if Fe3* is contaminated in the plating bath 12, the zinc, the iron ion and the pH does not become unbalanced because Fe3* is reduced on the gas diffusion electrode 1 to Fe21. Accordingly, the circulation of the plating solution of the plating bath 12 to an electrolytic reduction apparatus equipped with an ion exchange membrane is unnecessary, and the platting solution of the platting bath 12 can be effeclively used without being discharged. No variation of the thickness of the zinc-iron alloy layer plated on the steel 16 and no partial roughness of the surface is observed

Since the gas diffusion electrodes 1 functioning as an anode are supported in the reinforcing frame 4, providing a high mechanical strength, the electrodes are not flapped nor bent even when the steel 16 moves in the small plating bath 12 at a speed as high as 200 m/mln. The protection member 5 disposed on the front surface of the reinforcing frame 4, prevents the moving steel 16 from contacting the gas diffusion electrode 1 so that scratches and sparks are not produced.

The invention will now be illustrated by Examples which, however, are to be considered a merely exemples y practice of the invention, and not as a fimilation.

Example

A gas diffusion electrode 1 having a length of 100 mm and a width of 100 mm was formed by a gas diffusion layer 2 with a thickness of 500 µm and a reaction layer 3 with a thickness of 100 $\mu m_{\rm c}$ attached together. The gas diffision layer 2 consisted of hydrophobic carbon black of average particle size 420 Å and polytetrafluoroethylene of average particle size 0.3 µm. The reaction layer 3 consisted of hydrophilic carbon black of average particle size 420 Å, carrying a platinum catalyst load of 0.56mg/cm², hydrophobic carbon black of average particle size 420 Å and polytetrafluoroethylene of average particle size 0.3 µm. Ten gas diffusion electrodes 1 were installed and supported in an electroconductive reinforcing frame 4 made of titanium. A protection member 5 was equipped on the front surface of the reinforcing frame 4 at

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the reaction layer 3 side of the gas diffusion electrode f.

As shown in Fig. 2, a gas diffusion electrode apparatus 9 was disposed in an electrolytic cell 10, and a chloride bath 12 consisting of 50g/liter-FeCl₂ . nH₂ O. 245g/liter-ZnCl₂ . 1.5g/liter-NaPH₂O₂ (pH=3.0) was filled in the cell chamber 11 formed on the reac-Hon layer 3 side of the gas diffusion electrode 1. A hydrogen gas was successively introduced into the cell chamber 13, on the gas diffusion layer 2 side of the gas diffusion electrode 1. At the same time, current was flown to the respective gas diffusion electrodes 1 through the latticed reinforcing frame 4. Under these circumstances, a steel cathode 16 was passed through the plating bath 12 of the cell chamber 11 at a speed of 200m/min., at a current density of 150 A/drm² and a bath voltage of 8.6 V so that the steel 16 was electroplated with a zinc-iron alloy layer having a thickness of 4 µm.

The gas diffusion electrode 1 was employed as an anode while supplying hydrogen gas to the gas diffusion layer 2 side of the gas diffusion electrode 1 so as to make the anode reaction of the chloride platting bath 12 to be a hydrogen oxidation reaction. No chlorine gas was evolved on the anode. The zinc-iron alloy layer plated on the steel 16 was examined to have a smooth and gorgeous surface without burns. When the steel was successively passed to effect the electroplating of the zino-iron alloy, no iron ion was oxidized on the anode from Fe21 to Fe31. Even if Fe31 was contaminated in the platting bath 12, the zinc, the iron ion and the pH did not become unbalanced because Fe3+ was reduced on the gas diffusion electrode 1 to Fe21. Accordingly, the circulation of the plating solution of the plating bath an 12 to an electrolytic reduction apparatus equipped with the ion exchange membrane was unnecessary, and the plating solution of the platting bath 12 could be effectively employed without being discharged. No variation of the thickness of the zinc-tron alloy layer plated on the steel 16 and no partial roughness on the surface were ob-Served.

Stince the gas diffusion electrodes 1 functioning as an anode were supported in the reinforcing frame 4 so as to elevate the mechanical strength, the electrodes were not flapped nor bent even when the steel 16 moved in the narrow plating bath 12 at a speed as high as 200 m/min. Since the protection member 5 was formed on the front surface of the reinforcing frame 4, the moving steel 16 was never in contact with the gas permeable electrode 1 so that scratches and sparks were not produced.

Similar results were obtained when a suifale bath was used in place of the chlorine bath.

Clains

 A process for electroptating a zinc alloy on a steel cathode rapidly moving through a zinc alloy plating bath, in a cell equipped with a gas diffusion anode in the form of laminated film composed of a reaction layer and a gas diffusion layer bonded together, characterized by comprising

supporting said gas diffusion anode on a latticed electroconductive reinforcing frame and distributing current to said gas diffusion anode through said reinforcing frame:

preventing a sliding contact of said rapidly moving cathode with an opposing surface of said gas diffusion anode while restraining flapping of said gas diffusion anode by a protective latticed member disposed over said gas diffusion anode and fastened to said supporting reinforcing frame.

 A process as defined in claim 1, wherein said plating bath is a chloride bath and said steel cathode moves through the bath, at a speed as high as 200 m/min

Patentansprüche

 Verfahren zum Elektroplattieren einer Zinklegierungrung auf eine sich rasch durch ein Zinklegierung-Plattierungsbad bewegende Stahlkathode in einer Zelle, die ausgestattet ist mit einer Gasdiffusionsanode in Gestatt einer aus einer Reaktionsschlicht und einer Gasdiffusionsschicht, die eneinander befestigt sind, bestehenden laminierten Schicht,

dadurch gekennzeichnet,

daß es aufweist

Tragen der Gasdiffusionsanode au einem gitterförmigen, elektrisch leitfähigen, verstärkenden Rahmen und Verteilen von Strom auf die Gasdiffusionsanode durch den verstärkenden Rehmen;

Verhindern einer gleitenden Berührung der sich schnell bewegenden Kathode mit einer gegenüberliegenden Oberfläche der Gasdiffusionsande unter Eingreuzung eines Hin- und Herbewegens der Gasdiffusionsanode durch ein schülzendes, gitterförmiges Element, das über der Gasdiffusionsanode angeordnet und an dem tragenden, verstörkenden Rahmen befestigt ist.

 Verfahren nach Anspruch 1, bei dem das Plattierungsbad ein Chlorid-Bad ist und sich die Stahlkathode mit einer Geschwindigkeit bis zu 200 m/min durch das Bad bewegt.

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Revendications

- 1. Procédé de dépôt électrolytique d'un alliage de zinc sur une cathode en acier se déplaçant rapidement à travers un bain de dépôt électrolytique d'alliege de zinc dans une cellule munte d'une anode à diffusion gazeuse ayant la forme d'un film stratifié fait d'une couche de réaction et d'une couche de ditfusion des gaz liées ensemble, caractérisé en ce qu'il comprend les étapes consistant à:
 - supporter ladite anode à diffusion gazeuse dans un cadre de renfort électroconducteur à mailles et envoyer un courant à ladite anode à diffusion gazeuse par l'intermédiaiare dudit cadre de renfort,
 - empécher la contact glissant entre ladite cethode qui se déplace rapidement et une surface en vis-à-vis de ladite anode à diffusion gazeuse tout en évitant un pliage de ladite anode à diffusion gazeuse grâce à un elément de protection à mailles placé audessus de ladite anode à diffusion gazeuse et fixé audit cadre de remort et de soutien. 26
- 2. Procédé selon la revendication 1, dans lequel ledit bain de dépôt électrolytique est un bain de chlorures et ladite cathode en acler se déplace dans le bain à une vitesse aussi élevée que 200 m/min.

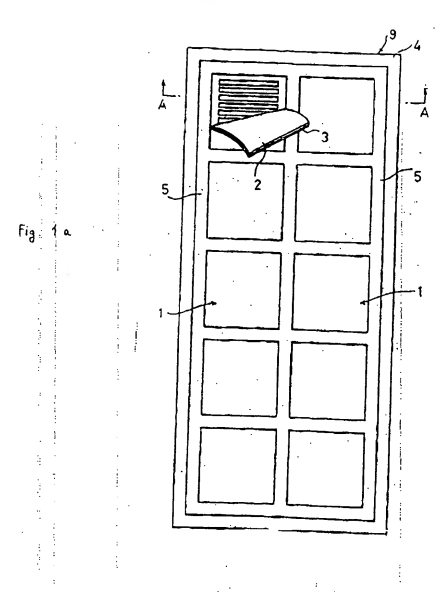


Fig. 1 b

